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Functional trait dissimilarity drives both species complementarity and competitive disparity

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Abstract: 1. Niche complementarity and competitive disparity are driving mechanisms behind plant community assembly and productivity. Consequently, there is great interest in predicting species complementarity and their competitive differences from their functional traits as dissimilar species may compete less and result in more complete use of resources. 2. Here we assessed the role of trait dissimilarities for species complementarity and competitive disparities within an experimental gradient of plant species richness and functional trait dissimilarity. Communities were assembled using three pools of grass and forb species based on a priori knowledge of traits related to (1) above- and below-ground spatial differences in resource acquisition, (2) phenological differences or (3) both. Complementarity and competitive disparities were assessed by partitioning the overyielding in mixed species communities into species complementarity and dominance effects. 3. Community overyielding and the underlying complementarity and competitive dominance varied strongly among the three plant species pools. Overyielding and complementarity were greatest among species that were assembled based on their variation in both spatial and phenological traits. Competitive dominance was greatest when species were assembled based on spatial resource acquisition traits alone. 4. In communities that were assembled based on species variation in only spatial or phenological traits, greater competitive dominance was predicted by greater differences in SLA and flowering initiation respectively, while greater complementarity was predicted by greater dissimilarity in leaf area and flowering senescence respectively. Greater differences in leaf area could also be linked to greater species complementarity in communities assembled based on variation in both phenological and spatial traits, but trait dissimilarity was unrelated to competitive dominance in these communities. 5. Our results indicate that complementarity and competitive disparity among species are both driven by trait dissimilarities. However, the identity of the traits that drives the complementarity and competitive disparity depends on the trait variation among species that comprise the community. Moreover, we demonstrate that communities assembled with the greater variation in both spatial and phenological traits show the greatest complementarity among species.

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**Functional trait dissimilarity drives both species
complementarity and competitive disparity**

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32 **Summary**

33 **1.** Niche complementarity and competitive disparity are driving mechanisms behind
34 plant community assembly and productivity. Consequently, there is great interest in
35 predicting species complementarity and their competitive differences from their
36 functional traits as dissimilar species may compete less and result in more complete
37 use of resources.

38 **2.** Here we assessed the role of trait dissimilarities on species complementarity and
39 competitive disparities within an experimental gradient of plant species richness and
40 functional trait dissimilarity. Communities were assembled using three pools of grass
41 and forb species based on a priori knowledge of traits related to (1) above- and
42 belowground spatial differences in resource acquisition, (2) phenological differences,
43 or (3) both. Complementarity and competitive disparities were assessed by
44 partitioning the overyielding in mixed species communities into species
45 complementarity and dominance effects.

46 **3.** Community overyielding and the underlying complementarity and competitive
47 dominance varied strongly among the three plant species pools. Overyielding and
48 complementarity was greatest among species that were assembled based on their
49 variation in both spatial and phenological traits. Competitive dominance was greatest
50 when species were assembled based on spatial resource-acquisition traits alone.

51 **4.** In communities that were assembled based on species variation in only spatial or
52 phenological traits greater competitive dominance was predicted by greater
53 differences SLA and flowering initiation respectively, while greater complementarity
54 was predicted by greater dissimilarity in leaf area and flowering senescence,
55 respectively. Greater differences in leaf area could also be linked to greater species
56 complementarity in communities assembled based on variation in both phenological

57 and spatial traits, but trait dissimilarity was unrelated to competitive dominance in
58 these communities.

59 **5.** Our results indicate that complementarity and competitive disparity among species
60 are both driven by trait dissimilarities. However, the identity of the traits that drives
61 the complementarity and competitive disparity depends on the trait variation among
62 species that comprise the community. Moreover, we demonstrate that communities
63 assembled with the greater variation in both spatial and phenological traits show the
64 greatest complementarity among species.

65

66 **Key-words** biodiversity, competition, community ecology, Jena Experiment, Trait
67 Based Experiment (TBE)

68

69

70 **Introduction**

71 It has long been observed that plant species mixtures are frequently more
72 productive than the average of the respective species monocultures; often referred to
73 as an overyielding effect (Darwin 1859; de Wit 1960; Hector et al. 1999; Tilman et al.
74 2001). However, the mechanisms by which diversity drives overyielding are still
75 widely debated as species diversity-productivity relationships are highly variable
76 (Adler et al. 2011). Furthermore, species number alone holds little information as to
77 how species interact and function as a community to drive the overall functioning of a
78 diverse community of species (Petchey & Gaston 2002). Thus, functional traits that
79 reflect the species strategies for acquiring resources are considered to be a key to
80 empirically assess, and predict, how species partition and compete for the local
81 resource pool that drives the productivity in mixed species communities (Chesson
82 2000; Lavorel & Garnier 2002; Ackerly & Cornwell 2007).

83 Conceptual frameworks emphasizing the importance of functional traits to
84 infer niche differences among species are based on the hypothesis that phenotypic and
85 phylogenetically similar species have similar ecological requirements and will thus
86 compete more strongly; coined the ‘limiting-similarity hypothesis’ (MacArthur &
87 Levins 1967; Mayfield & Levine 2010). Under the trait-similarity hypothesis, it can
88 be expected that greater differences among species in functional traits should reflect
89 greater niche differentiation that allows species to avoid competition and partition the
90 local resource pool (Westoby et al. 2002; Falster & Westoby 2003; Grime 2006;
91 Cadotte et al. 2013; Kraft et al. 2014). For instance, resource partitioning may arise
92 from differences in plant height, rooting patterns, and phenology that allow different
93 species to utilize different spatial and temporal resources. Although taller or more
94 shallow rooted plant species may have priority in acquiring particular resources, they

95 are likely not able to completely capture all available resources due to trade-offs in
96 optimizing certain resource capture traits over others (Westoby et al. 2002; Falster &
97 Westoby 2003; Wright et al. 2004; Reich 2014; Diaz et al. 2016). This provides
98 opportunity for species with a different suite of traits optimized for capturing the
99 resource margin, such as those species specialized in capturing understory light or
100 resources deeper in the soil profile, to coexist and contribute to community-level
101 resource capture and overyielding (Liira & Zobel 2000; Fargione & Tilman 2005;
102 Aarssen et al. 2006). Greater resource use partitioning in mixed species communities
103 is often considered as niche ‘complementarity’ that can be reflected in the greater
104 performance of species in mixture relative to their monoculture performance (Loreau
105 & Hector 2001; Hector et al. 2002).

106 However, greater differences among species in functional traits may also
107 reflect differences in their competitive abilities to capture resources over competing
108 neighbouring species. For instance, taller and larger plants can pre-empt light over
109 smaller plants (Freckleton & Watkinson 2001; Falster & Westoby 2003; Weiner &
110 Damgaard 2006; Roscher et al. 2015). Belowground, greater root-length density can
111 provide a species with a competitive advantage in capturing more soil nutrients over
112 neighbouring species (Casper et al. 1997; Fargione & Tilman 2006; Fort et al. 2014).
113 Therefore, species with a particular suite of traits that are more favourable in pre-
114 emptying resource capture within a given environment, relative to neighbouring
115 species, can result in a competitive hierarchy among species; referred to as the
116 ‘competition-trait hierarchy’ hypothesis (Mayfield & Levine 2010; Kunstler et al.
117 2012; Kunstler et al. 2016). Plants with traits that allow them to effectively acquire
118 the local resources that provides them with a better competitive advantage over
119 neighbouring plant species can result in the species disproportionate contribution to

120 the productivity in a mixed species community. The presence of a particularly
121 productive species that drives the productivity in a community can be considered as a
122 ‘selection’ effect, where the inclusion of a particularly productive species drives the
123 community performance (Loreau 2000; Loreau & Hector 2001). Further, a
124 ‘dominance effect’ can occur as part of the ‘selection effect when such highly
125 productive species drive the performance of the community at the expense of
126 subordinate species (Fox 2005).

127 Although functional trait differences among species can result in potential
128 opposing outcomes of competition under the ‘limiting-similarity’ and ‘competition-
129 trait hierarchy’ hypotheses (Mayfield & Levine 2010), both mechanisms may
130 influence community overyielding in diverse communities. Conveniently,
131 overyielding in plant communities can be partitioned into the ‘complementarity’
132 effect and the ‘selection’ effect (Hector & Loreau 2001). Both complementarity and
133 selection effects can operate simultaneously within a community (see Hector et al.
134 2002 for all potential scenarios) and sum to the overyielding of a community.
135 Importantly, the selection effect can be further partitioned into a ‘dominance’ effect
136 where particularly productive species result in reduced productivity of subordinate
137 species, as well as a ‘trait-complementarity’ effect where highly productive species
138 drive the community performance, but at no cost to subordinate species (Fox 2005).
139 However, there is little evidence as to how functional trait dissimilarities among
140 species relate to the complementarity, selection, and dominance effects that together
141 determine the overyielding effect of mixed species assemblages.

142 Here we test the hypotheses that the assembly of species with greater
143 functional trait dissimilarity will influence the overyielding of a community by
144 favouring either (i) niche partitioning and promote a complementarity effect, or (ii)

145 provide a competitive advantage over neighbouring species to promote a selection and
146 dominance effect. However, it is conceivable that the role of plant functional trait
147 dissimilarity among species within a community may depend upon the variation in
148 particular functional traits of species that comprise a community. Therefore, we assess
149 the conditionality of our hypotheses that increased trait dissimilarity among species
150 predicts complementarity and competitive interactions by using three different pools
151 of plant species that were known to vary in 1) spatial above- and belowground traits
152 (pool 'S'), 2) phenological traits (pool 'P'), or 3) both spatial and phenological traits
153 (pool 'SP'). We tested our hypotheses in experimental grassland plant communities
154 that were designed to represent a gradient of plant species richness crossed with
155 varying levels of pre-determined functional trait diversity levels within each of the
156 three species pools (Ebeling et al. 2014). Such an experimental design allows us to
157 explore the influence of spatial and temporal functional trait compositions on the
158 underlying competitive and complementarity mechanisms that determine the overall
159 functioning of a community.

160

161 **Materials and methods**

162 *Experimental design*

163 In order to test our hypotheses that greater trait differences among species can
164 predict niche complementarity or competitive disparity we used 20 plant species (see
165 Table S1 in Supporting Information) selected from the pool of 60 species that occur
166 within the Jena Experiment (Roscher et al. 2004). These 20 plant species were
167 selected for their variation in their phenology as well as aboveground and
168 belowground resource-acquisition traits (Ebeling et al. 2014). Specifically, plant
169 height, leaf area, rooting depth, and root length density were considered as traits

170 associated with spatial resource acquisition, while the date of growth and flowering
171 initiation were considered as phenological traits. Based on these traits, plant species
172 were grouped into three partially overlapping pools of species that vary in trait
173 dissimilarities from being highly similar to highly dissimilar (see Ebeling et al. 2014
174 for details). In brief, species trait variation was assessed by PCA, where the PCA axis
175 1 separated species based upon their above-belowground spatial resource use traits,
176 while PCA axis 2 separated species based on phenological traits (see Fig. S1). Species
177 in pool ‘S’ species were selected based on variation in above- and belowground
178 spatial traits (i.e. selected across a range of small to large differences along the PCA
179 axis 1). Species in pool ‘P’ were selected based on variation in phenology (i.e.
180 selected across a range of small to large differences along the PCA axis 2). Pool ‘SP’
181 consisted of species that were selected for their variation in both spatial and
182 phenological traits. Each pool consisted of eight species with different combinations
183 of grasses, tall forbs, and small forbs according to the functional group classification
184 of the Jena Experiment (Roscher et al. 2004). Four of the 20 species occur in two
185 species pools (see Table S1). Further details on the selection of species are described
186 in detail in Ebeling et al. (2014)

187 The experimental plant communities were sown into 138 plots within the Jena
188 Experiment field site located near Jena, Germany (50° 57’ 3” N, 11° 37’ 35” E, 130 m
189 a.s.l.). Initial sowing took place in autumn 2010 and re-sown in summer 2011 ensure a
190 successful establishment of the plant communities and all sown plant species were
191 observed to be present in the plots during data collection in 2012. Plots were 3.5 × 3.5
192 m in size and arranged in three spatial blocks to account for edaphic factors along the
193 Saale River. Plant communities were sown in plots with a species richness of 1, 2, 3,
194 4, or 8 plant species from a given species pool, and the species combinations in each

195 plot covered a range of functional trait dissimilarities (Ebeling et al. 2014). Species
196 were sown in equal proportions with a total density of 1000 germinable seeds per m⁻²
197 based on laboratory germination rates, such that the expected initial number of
198 individuals per species would be relatively equivalent. In 2011 and 2012, plots were
199 weeded in summer (July) and autumn (October), as well as in early spring (March)
200 2012, in order to maintain the sown plant community composition. Biomass harvest
201 occurred in 2012 in spring (late May) and was harvested again in summer (late
202 August). The biomass of each plot was harvested by cutting all plants approximately 3
203 cm above the soil surface within two randomly allocated 0.1 m² subplots within each
204 plot, which is typical in long-term plant biodiversity experiments with large plot size
205 (Tilman et al. 1997; Hector et al. 1999; Tilman et al. 2001; Roscher et al. 2004;
206 Marquard et al. 2009; Roscher et al. 2011). Following each harvest, the plots were
207 mown to a height of 5 cm as is typical for managed hay meadows in the region.
208 Harvested plant material was then sorted to species. Biomass of species that were not
209 initially sown into the plots was pooled together as weed biomass. All plant material
210 was dried at 70°C for 48 h and subsequently weighed to quantify the biomass of
211 individual sown plant species and weed biomass. The spring and summer biomass
212 measurements of each species were summed to obtain the annual productivity.

213

214 *Species traits*

215 In 2012, we collected trait data on the 20 plant species. Traits were measured
216 in the species monocultures of the same year in which the biomass data was collected
217 so that the measured trait values for each species coincides with the observed biomass
218 production of each species. We measured flowering initiation and cessation as
219 phenological traits that reflect the seasonal timing in resource capture as well as

220 above- and belowground traits that reflect differences in how resources are captured
221 spatially (Table 1). Flowering initiation was quantified as the week of the year in
222 which the first flowers of a species appeared. Flowering cessation is the week of the
223 year in which all flowers of the species have senesced. The maximum canopy height
224 was measured just prior to harvesting biomass by averaging five canopy height
225 measurements along a transect across the plot. Leaf characteristics were measured by
226 sampling a total of 5-12 fully developed leaves from different individuals. Leaves
227 were stored in moist paper in sealed plastic bags at 4°C overnight for rehydration and
228 then blotted dry to remove any surface water. Leaf fresh weight was then measured,
229 and the leaf surface area was measured with a portable Leaf Area Meter (LI-3100,
230 LICOR, Lincoln, USA). Afterwards, leaf samples were dried at 70°C (48 h), and dry
231 weights were recorded. Specific leaf area (SLA) was calculated as the ratio of leaf
232 area to dry weight ($\text{mm}^2 \text{mg}^{-1}$), and leaf dry matter content (LDMC) is the ratio of dry
233 weight to fresh weight (mg g^{-1}). Leaf area (cm^2) was calculated as the average of the
234 sampled leaves.

235 Root traits were determined by taking eight soil cores (4 cm in diameter and
236 40 cm in depth) that were sectioned into depths of 0-5, 5-10, 10-20 cm, 20-30, and 30-
237 40 cm (Ravenek et al. 2014). Sections were pooled by depths per plot and washed
238 clean of soil and roots less than 2 mm diameter were stored in 70% EtOH. A
239 subsample of roots was stained with neutral red mixed in 70% EtOH overnight and
240 the root length was determined by scanning stained roots (600 dpi, Epson Expression
241 10000 XL scanner, Regent Instruments, Quebec, Canada) using WinRhizo software
242 (Regent Instruments; manual pixel classification: 225). All root samples were then
243 dried at 65°C for at least 48 h and weighed. Specific root length (SRL) was calculated
244 as the ratio of root length to dry mass of the subsample ($\text{mm}^2 \text{mg}^{-1}$). Mean rooting

depth (MRD) was calculated for each plot as the weighted mean of root mass per layer. Root mass density (RMD) was calculated from root mass per cm^3 of soil and averaged for each layer. Root length density (RLD) was calculated as the root mass per root length (mg cm^{-1}) and averaged for each layer. Due to the time and resource constraints on sampling and processing required to quantify root traits, root traits were only measured in the monocultures of species pools 'S' and 'P'.

251

252 *Community level indices*

In order to test for community wide complementarity and competitive disparity among species we calculated the complementarity and selection effects were calculated following Loreau & Hector (2001) using the annual biomass production of the species. Specifically, we first calculated species relative yields (RY); which is the observed biomass of species in a mixture (O) divided by the species monoculture (M) such that $RY = O/M$ (de Wit 1960). The complementarity effect is calculated as $N\overline{\Delta RY}\overline{M}$, where N is the number of species in the mixture, ΔRY is the difference in the RY of a species from its expected relative yield ($1/N$) and M is the monoculture biomass. The complementarity effect is positive when species perform better than expected in the mixture on average and negative when performing poorly in mixture. The selection effect is calculated as $N\text{cov}(\Delta RY, M)$ and is positive when highly productive species in monoculture are also highly productive in mixtures and negative when highly productive species contribute less than expected in the mixture. To better assess competitive disparity within the communities we further partitioned the selection effect into the dominance effect, calculated as $N\text{cov}(M, RY/\Sigma RY - 1/N)$ and the trait-complementarity effect, calculated as $N\text{cov}(RY - RY/\Sigma RY, M)$, following Fox (2005). The dominance effect is positive when highly productive species dominate the

270 mixture at the expense of less productive species and negative when less productive
271 species dominate the mixture at the expense of more productive species. The trait-
272 complementarity effect is positive when highly productive species drive the
273 productivity of the mixture, but not at the expense of less productive species and
274 negative when less productive species drive the productivity of the mixture, but with
275 no negative effect on the more productive species.

276 The ten plant traits were standardized ($\mu = 0$, $\sigma = 1$) and used to calculate
277 Rao's quadratic entropy (*RaoQ*, un-weighted by species abundance) as an index of
278 functional trait dissimilarity. We used *RaoQ* since it was also used in the initial
279 experimental design (Ebeling et al. 2014), and it is highly similar to functional trait
280 dispersion '*FDis*' (Laliberté & Legendre 2010; Clark et al. 2012). *RaoQ* was
281 calculated separately for each species pool using all 10 traits, or only phenology,
282 aboveground or belowground traits separately in order to identify the effect of the
283 different suite of traits (aboveground, belowground or phenology). The *RaoQ* using
284 all 10 traits was standardized (mean = 0, SD = 1) by species pool since not all traits
285 were measured in pool 'SP' (i.e. no belowground traits measured). The mean pairwise
286 difference (MPD) in individual traits was also calculated in order to assess the
287 importance of the difference among species in individual traits.

288

289 *Data analyses*

290 All data processing and analyses were carried out using R version 3.02 (R
291 Core Team, 2014). We first tested for overall effects of species pool, sown plant
292 species richness, and the overall trait dissimilarity on the productivity of the
293 communities, by ANOVA with species pool, richness, trait dissimilarity (*RaoQ* using
294 all traits), and the interactions between pool with richness and trait dissimilarity as

295 main terms. Block was included as a covariate. A contrast term testing for differences
 296 between the productivity of mixtures and monocultures (overyielding effect) and its
 297 interaction with species pool was also included. To test the hypothesis that greater
 298 trait dissimilarity can predict complementarity and competitive disparities we
 299 assessed the overyielding, complementarity, selection, dominance, and trait-
 300 complementarity effects as above, but omitting the contrast between monocultures
 301 and mixtures (since effects are only calculated in mixtures). Secondly, we
 302 independently assessed the effect of dissimilarity in phenological, aboveground, and
 303 belowground traits among species by substituting the trait dissimilarity (all traits)
 304 term in the above ANOVAs with the trait dissimilarity in only phenological,
 305 aboveground, or belowground traits. Finally, individual functional trait differences
 306 (calculated as MPD) were then assessed for their ability to predict complementarity,
 307 selection, dominance, and trait-complementarity effects by first standardizing ($\mu = 0$,
 308 $\sigma = 1$) all dependent variables (complementarity, selection, dominance, and trait-
 309 complementarity) and independent variables (all MPD indices) so that parameter
 310 estimates were comparable on a common scale. The standardized data were then used
 311 in lasso regression (Least Absolute Shrinkage and Selection Operator) for generalized
 312 linear model selection with penalized maximum likelihood (Tibshirani 1996,
 313 Friedman et al. 2010). The shrinkage parameter (λ) was determined using 10-fold
 314 cross-validation to obtain a λ that provided the minimum mean squared error using
 315 the function 'cv.glmnet' in the R package 'glmnet'. Terms with non-zero coefficients
 316 were then included in a single linear multiple regression model.

317

318 **Results**

319 *Relationships between species richness, trait dissimilarity and productivity*

320 We found that the species pool had the strongest effect on the overall
 321 productivity, where species pool ‘SP’ (species varying in both spatial and
 322 phenological traits) was significantly less productive than pools ‘S’ (species varying
 323 more in spatial traits) and ‘P’ (species varying more in phenological traits) (Fig. 1a,
 324 $F_{2, 124} = 12.93$, $P < 0.001$, Table S2). Species richness had little overall effect on the
 325 net productivity of the communities (Fig. 1a, $F_{1, 124} = 0.37$, $P = 0.542$) and showed no
 326 interaction with species pool ($F_{2, 124} = 0.22$, $P = 0.806$, Table S2). There was an
 327 overall overyielding effect (mixed species communities performing better than the
 328 monoculture average, $F_{2, 124} = 4.70$, $P = 0.032$, Table S2), which did not differ
 329 significantly among species pools ($F_{2, 124} = 1.57$, $P = 0.212$). Moreover, we found
 330 productivity to be positively related to greater trait dissimilarity depending on the
 331 species pool (Fig. 1b, $F_{2, 124} = 3.86$, $P = 0.024$, Table S2), where the trait
 332 dissimilarity-productivity relationship was only significant in species pool ‘SP’.

333

334 *Effects of trait dissimilarity on overyielding, complementarity and selection effects*

335 Overyielding, complementarity and the selection effects were not significantly
 336 related to plant species richness (overyielding: $F_{1, 103} = 0.69$, $P = 0.408$,
 337 complementarity: $F_{1, 103} = 0.50$, $P = 0.482$, selection: $F_{1, 103} = 0.13$, $P = 0.718$, see
 338 Table S3). However, both overyielding and complementarity were influenced by
 339 greater overall trait dissimilarity depending on the species pool (overyielding: $F_{2, 103} =$
 340 4.83 , $P = 0.010$, complementarity: $F_{1, 103} = 5.11$, $P = 0.008$). More specifically, we
 341 found the complementarity effect in species pool ‘P’, declined with greater
 342 dissimilarity in phenological traits (Fig. 2a). However, complementarity was not
 343 related to phenological dissimilarity in species pools ‘S’ and ‘SP’, resulting in a
 344 marginal interaction between species pool and phenological trait dissimilarity ($F_{1, 103}$

345 = 2.80, $P = 0.066$, Table S3). Greater dissimilarity in phenological traits had little
 346 effect on overyielding in all three species pools (Fig. 2a, Table S3). The selection
 347 effect was also unrelated to phenological dissimilarity among species in each of the
 348 three species pools (Fig. 2a).

349 Greater dissimilarity in aboveground traits had the strongest effect on
 350 increasing overyielding and the selection effect in species pool ‘S’ (Fig. 2b).

351 However, these relationships did not differ significantly from the same relationships
 352 in the other two species pools (overyielding: $F_{1, 103} = 1.88$, $P = 0.158$, selection: $F_{1, 103}$
 353 $= 1.57$, $P = 0.212$, Table S3). Increasing aboveground trait dissimilarity in pool ‘S’
 354 was not to complementarity, indicating that the positive effect of aboveground trait
 355 dissimilarity on overyielding was due to an increased selection effect (Fig. 1b).

356 Aboveground trait dissimilarity had no effect on the overyielding, complementarity
 357 and selection effects in species pool ‘P’. However, in species pool ‘SP’, both
 358 complementarity and overyielding increased with greater dissimilarity among species
 359 in aboveground traits, but not the selection effect (Fig. 1b), illustrating that the
 360 overyielding in species pool ‘SP’ was driven by a complementarity effect. The strong
 361 effect of aboveground trait dissimilarity on the complementarity effect in these
 362 communities (pool ‘SP’) resulted in a significant interaction effect between species
 363 pool and aboveground trait dissimilarity ($F_{1, 103} = 3.87$, $P = 0.024$, Table S3). The
 364 dissimilarity in belowground traits had little overall effect on the overyielding,
 365 complementarity and selection effects (Fig. 2c, Table S3). Overall, the species pool
 366 had the strongest effect on the overyielding, complementarity and selection effects
 367 (Fig. 2d, Table S3). Overyielding and the complementarity effect were lowest in
 368 species pool ‘S’ followed by species pool ‘P’ and were greatest in species pool ‘SP’.

369 Conversely, the selection effect was greatest in species pool ‘S’ and lowest in species
370 pool ‘SP’ (Fig. 2d).

371

372 *Effects of trait dissimilarity on the dominance effect*

373 By partitioning the selection effect into the “trait-complementarity” and
374 “dominance” effect (Fox 2005), we found that both the dominance and trait-
375 complementarity effects were greater in species pool ‘S’ and ‘P’ than in species pool
376 ‘SP’, as observed in the selection effect (Fig. 3, Table S4, and see Fig. S2). However,
377 the trait-complementarity effect was unrelated to phenological, aboveground or
378 belowground trait dissimilarity (Table S4). The dominance effect was only
379 significantly and positively related to the aboveground trait dissimilarity in species
380 pool ‘S’ resulting in a significant species pool by aboveground trait dissimilarity
381 interaction effect (Fig. 3, $F_{1,103} = 7.91$, $P = 0.001$, Table S3).

382

383 *Identifying effects of specific trait differences*

384 Of the specific trait dissimilarities among species, we found greater
385 dissimilarities in SLA could best predict both the selection and dominance effects in
386 species pool ‘S’ (Fig. 4a). Additionally, we found that greater differences in leaf area
387 best predicted greater complementarity in pool ‘S’ (Fig. 4a). In pool ‘P’ the selection
388 and dominance effects were predicted by greater flowering initiation, while a lower
389 species complementarity was predicted by greater dissimilarity in flowering
390 senescence (Fig. 4b). The selection and dominance effects in communities from pool
391 ‘SP’ were poorly predicted by the dissimilarity in all traits, but greater differences in
392 leaf area best predicted greater species complementarity (Fig. 4c).

393

Discussion

Here we experimentally manipulated species functional trait dissimilarities to assess the role of species diversity and functional trait dissimilarities in predicting species complementarity and competitive disparities. Overall, we found that the pool from which species were assembled had the strongest influence on productivity, overyielding, and its underlying complementarity and competitive dominance effects. Specifically, the assembly of species chosen for the greatest variation in both spatial and temporal resource acquisition traits (species pool ‘SP’) resulted in the greatest complementarity, which drove overyielding in these communities. This shows that the assembly of communities composed of species that vary in both temporal and spatial resource acquisition traits can predict greater overyielding through greater complementarity, providing support for the competition-trait similarity hypothesis (*i*). However, our results also reveal that the assembly of species based on variation in only spatial traits resulted in the community overyielding to be driven by greater competitive dominance. This provides support for the second hypotheses (*ii*) that greater functional differences among competitors lead to competitive disparity. These results parallel findings that a greater difference in functional traits between competitors can result in a ‘competition-trait hierarchy’ (Freckleton & Watkinson 2001; Kunstler et al. 2012; Fort et al. 2014; Kunstler et al. 2016). Overall, our study demonstrates that competitive disparity among species can occur through spatial or temporal resource pre-emption when species vary most in spatial or temporal resource capture traits respectively (e.g. species pools ‘S’ and ‘P’), but also that species complementarity is best predicted when species vary more greatly in both temporal and spatial traits (e.g. species pool ‘SP’).

419 *Effects of trait differences on species complementarity*

420 The strong effect of the species pool in our study suggests that niche
421 complementarity may be best predicted by greater differences in phenology in
422 combination with greater differences in spatial resource acquisition, compared to only
423 considering variation in spatial or phenological resource acquisition strategies alone.
424 The finding that the combination of both spatial and temporal niche segregation, as
425 reflected in species functional trait differences, together play a key role in species
426 resource use complementarity is supported by previous findings that interactions
427 among plant species are temporally dynamic (Connolly et al. 1990; McKane et al.
428 1990; Fargione & Tilman 2005).

429 The assessment of individual trait differences revealed that greater differences
430 in leaf area could best predict greater complementarity within species pools ‘S’ and
431 ‘SP’, where the species spatial resources acquisition traits were varied. The link
432 between larger leaf area differences and a greater complementarity effect is likely
433 reflective of differences among species in light absorption strategies, such as investing
434 differently in fewer larger leaves versus more numerous smaller leaves (Milla &
435 Reich 2007; Whitman & Aarssen 2010). For instance, in our species pool ‘SP’, where
436 complementarity was greatest and best predicted by leaf area differences, was
437 composed mostly of small and tall forbs. Thus, the positive complementarity effect
438 that drove overyielding in these communities may reflect that these two growth forms
439 are adapted in leaf production strategies to minimize competition in light absorption
440 as reflected in their differences in leaf area (Falster & Westoby 2003; Aarssen et al.
441 2006; Whitman & Aarssen 2010; Reich 2014). Such differences in leaf production
442 strategies are thought to be a key mechanism by which plant species coexist and avoid
443 competition (Aarssen et al. 2006; Wacker et al. 2009, Whitman & Aarssen 2010).

444 Although not measured, root traits in our species pool ‘SP’ may have provided further
445 information as to the increase in complementarity in these communities. However,
446 since rooting traits had no apparent influence in species pools ‘S’ and ‘P’ it also likely
447 had little effect in pool ‘SP’. Additionally, it has been observed that rooting traits may
448 not always relate to aboveground productivity during initial community
449 establishment, but may become important in time as the communities establish
450 (Mommer et al. 2010; Ravenek et al. 2014).

451 It should be considered that since the complementarity effect in our study is
452 the measure of average species performances in mixtures relative to their
453 monocultures, intraspecific density dependence mechanisms might have also
454 governed community productivity and complementarity aside from interspecific
455 differences in resource acquisition traits (Schöb et al. 2015). Negative density-
456 dependence in plants can result from the increase in plant species-specific pathogens,
457 herbivores, and the depletion of key soil resources required by the species (Maron et
458 al. 2011; Kulmatiski et al. 2012). These negative density-dependence mechanisms
459 could potentially explain the positive species complementarity we observed that
460 would not be captured by interspecific trait differences quantified in our study.

461

462 *Effects of trait differences on competitive disparity*

463 The similar increase in the selection and dominance effect with greater trait
464 differences, such as in species pool ‘S’ with SLA and in species pool ‘P’ with
465 flowering initiation, reveals that the selection effect in these communities was due to a
466 dominance effect (where the presence of species that are particularly productive in
467 monoculture drive the community productivity in mixtures at the expense of the less
468 productive species). This result is supported by previous work that indicates the

469 importance of size asymmetry in shaping competition and composition in plant
470 communities (Weiner 1990; Freckleton & Watkinson 2001; DeMalach et al. 2016).
471 Greater SLA values are known to be associated with increased photosynthesis,
472 nutrient uptake, and rapid growth that consequently provides the species with a
473 competitive advantage in productive systems (Poorter & Remkes 1990; van der Werf
474 et al. 1993; Knops & Reinhart 2000; Fargione & Tilman 2002). In our study, this
475 indicates that the greater differences in SLA among species that drove the competitive
476 dominance effect in species pool ‘S’ favoured larger, faster growing species. We also
477 observed a positive effect of greater dissimilarity in the timing of flowering initiation
478 in our species pool ‘P’ (assembled for variation in phenological traits) on the
479 dominance effect and a negative relationship between complementarity and
480 dissimilarity in flowering senescence. These results may reflect that greater temporal
481 resource pre-emption can favour the earlier developing plant species.

482 Although larger and faster growing species may hold a competitive advantage
483 during early stages of community establishment, during later stages in more
484 established communities, competition may favour slower-growing species (van der
485 Werf et al. 1993). For instance, it is often observed that species complementarity
486 increases, and selection or dominance effects decrease, as plant communities develop
487 and undergo compositional re-assembly (Tilman et al. 2001; Marquard et al. 2009;
488 Reich et al. 2012). Therefore, our trait dissimilarity-driven dominance parallels the
489 concept that communities initially sown with high functional trait variation can result
490 in competitive disparity among species during the early stages of community
491 establishment that shifts the proportional abundances of species in favour of more
492 productive species (Loreau 2000). The increasing dominance effect in our plant
493 communities at higher trait dissimilarity may be best reflective of greater community-

494 wide competition aboveground following the initial assembly of the communities;
495 which is in line with the first years of many longer running biodiversity experiments
496 (Tilman et al. 2001; Marquard et al. 2009; Roscher et al. 2011; Reich et al. 2012;
497 Ravenek et al. 2014). However, it should also be noted that the same suits of traits
498 did not predict the dominance effect and complementarity across the three species
499 pools in our study, reflecting the importance of the differences in the trait variation
500 among species within pools. Therefore, the way in which particular assemblages of
501 species may compete can depend upon the trait variation among the larger pool of
502 species from which it originates.

503

504 *Conclusion*

505 Our results provide evidence that species with greater differences in both
506 spatial and temporal resource acquisition strategies can result in greater
507 complementarity (i.e. tall versus small forbs in our pool ‘SP’). However, our study
508 also demonstrates that the assembly of plant species in an experimental functional
509 trait diversity gradient can result in greater competitive disparity during early
510 establishment of the community, likely through greater spatial and temporal resource
511 pre-emption as evidenced in our species pools ‘S’ and ‘P’. These results parallel a
512 slowly growing literature base that demonstrates greater functional trait dissimilarities
513 among species does not always relate to species complementarity and reduced
514 resource competition (Cahill et al. 2008; Mommer et al. 2010; Kunstler et al. 2012;
515 Fort et al. 2014). In our study, the increase in the competitive disparity among species
516 with greater SLA in communities assembled with greater variation in spatial resource
517 capture traits, and flowering initiation in communities assembled with greater
518 variation in phenological traits, indicates the competitive advantage of larger and

519 faster-growing species following initial assembly and early establishment in
520 productive landscapes (Poorter & Remkes 1990; van der Werf et al. 1993; Knops &
521 Reinhart 2000; Fargione & Tilman 2002). Overall our study provides empirical
522 insights as to how functionally dissimilar species establish and compete to shape
523 community composition in a novel environment in support of previous conceptual and
524 empirical studies on how plant traits may predict community assembly processes and
525 functioning (Loreau 2000; Grime 2006; Mayfield & Levine 2010; HilleRisLambers et
526 al. 2012). If competitive disparity begets functional complementarity in the temporal
527 assembly process of species compositions, tracking the temporal change in the link
528 between functional dissimilarities and community level performance will be of key
529 importance. Such empirical observations are required to gain a better mechanistic and
530 predictive understanding of ecological linkages among functional traits, stable
531 competition-coexistence mechanisms, and the multi-functionality of diversity.

532

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539

540 **Data Accessibility**

541 Data are available at: doi: XXXXXXXXXXXX

542

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713

714 **Tables**

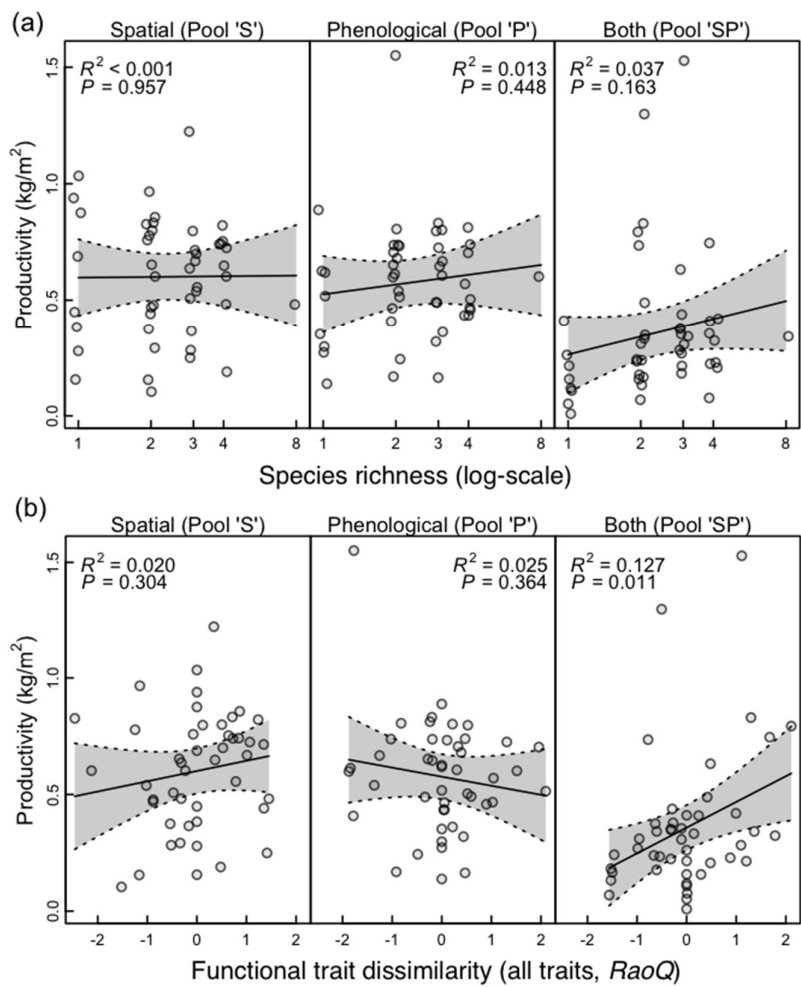
715 **Table 1.** List of traits that were measured on the species in their monocultures with
 716 abbreviations in parentheses and their measured units

Trait	Trait type	Units
1. Flowering initiation (F. init)	Phenological	week of year
2. Flowering senescence (F. sene)	Phenological	week of year
3. Specific leaf area (SLA)	Aboveground	mm ² /mg
4. Leaf dry matter content (LDMC)	Aboveground	mg/g
5. Leaf area (L. area)	Aboveground	cm ²
6. Maximum canopy height (Height)	Aboveground	cm
7. Mean Rooting Depth (MRD) †	Belowground	cm
8. Root Mass Density (RMD) †	Belowground	mg/cm ³
9. Specific Root Length (SRL) †	Belowground	mm/mg
10. Root Length Density (RLD) †	Belowground	mg/cm

717 † Traits only quantified in species pool ‘S’ and ‘P’ (see Methods)

718
 719

720 **Figures**
721



722
723 **Figure 1.** The relationship between the productivity in each plant species pool and (a)
724 the sown species richness and (b) the overall functional trait dissimilarity
725 (standardized by pool, see methods). Fit statistics (R^2 and P-values) are indicated for
726 each. Solid lines indicate regression relationships, and the grey shaded region is the
727 95% confidence band for the relationship.
728

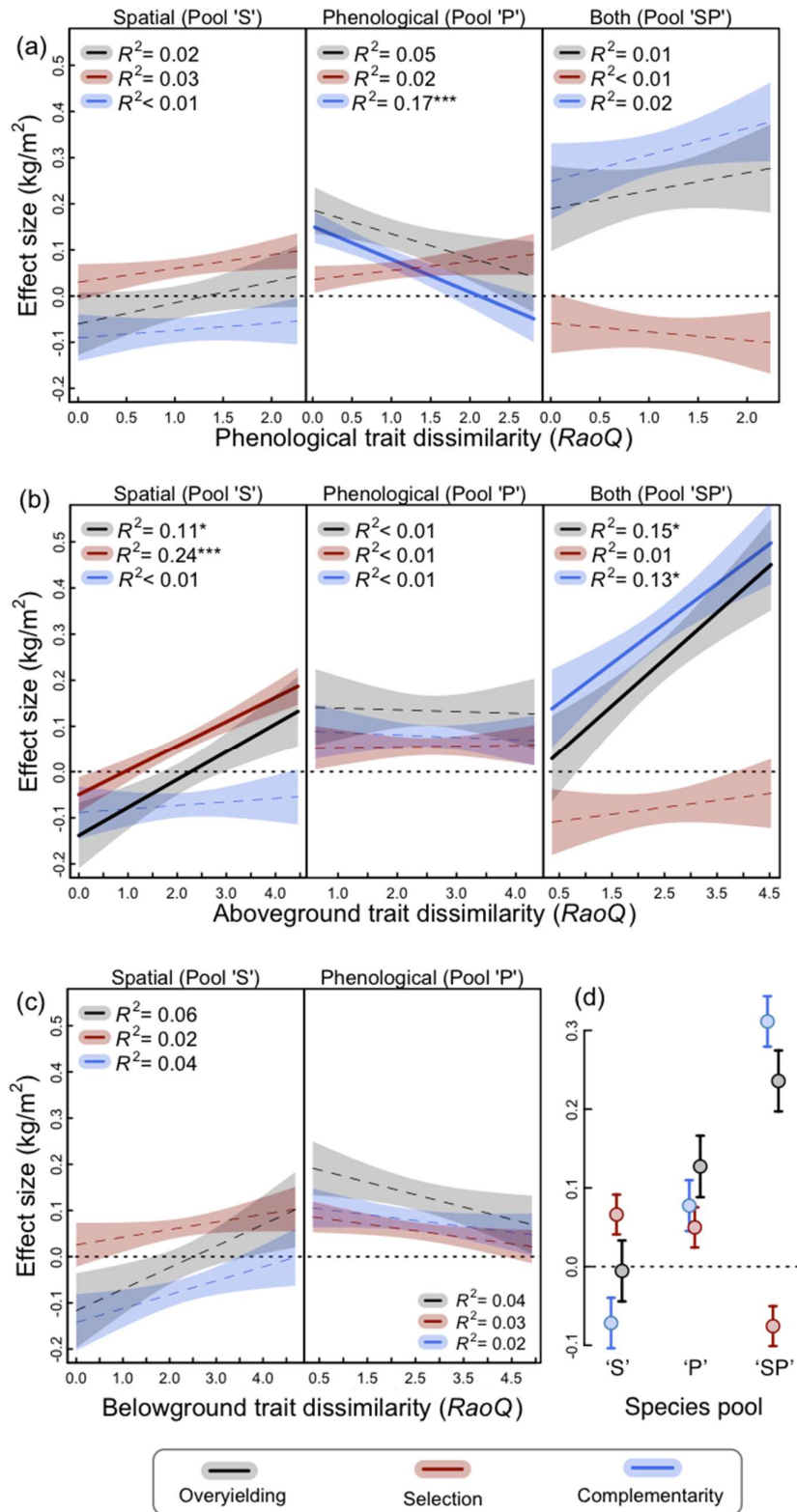
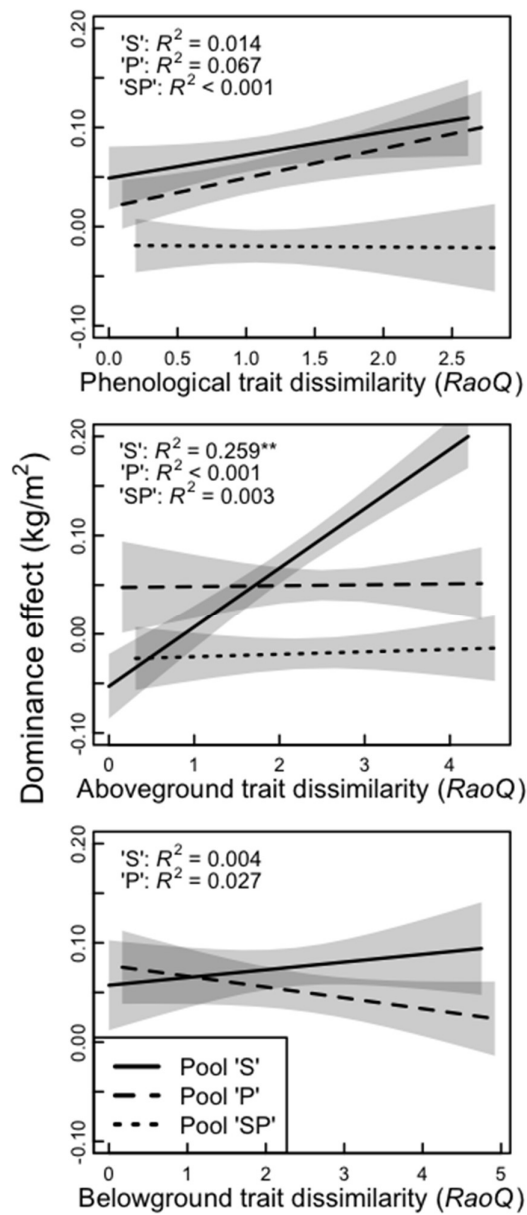


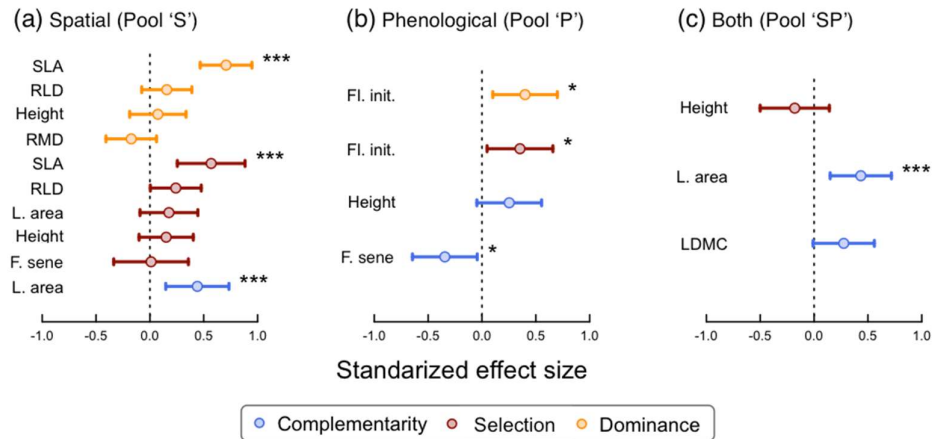
Figure 2. Relationships between the overyielding in species mixtures and the complementarity and selection effects with the variation in (a) phenological traits, (b) aboveground traits and (c) belowground traits (only measured in pool 'S' and 'P').

733 Solid lines indicate the regression relationships and shaded regions indicate the
734 standard error around the relationship. Fit statistics (R^2 and associated significance:
735 $*P < 0.05$, $***P < 0.001$) are indicated for each. Relationships are shown for each
736 species pool (shown in the panels from left to right). The overall mean overyielding,
737 selection and complementarity effects for each species pool are shown in (d) with
738 standard errors. Note the complementarity and selection effects sum to the
739 overyielding effect where greater overlap between overyielding with the selection or
740 complementarity effect indicates the greater the contribution of the selection or
741 complementarity effect to the overyielding of the community.
742



744

745 **Figure 3.** Relationships between the dominance effect and dissimilarity in
746 phenological traits, aboveground traits and belowground traits are shown for each
747 species pool (indicated by different regression lines). Lines indicate regression
748 relationships and the grey shaded regions are the standard errors for the regression fit.
749 The fit statistics (R^2) are provided for each ($^{**}P > 0.01$).



750

751 **Figure 4.** Standardized effects of the MPD in individual plant functional traits (see
 752 Table 1 for trait abbreviations) on the complementarity effect, selection effect, and the
 753 dominance effect. Error bars are 95% confidence intervals for a difference from 0 (no
 754 effect). Significance is indicated by: * $P < 0.05$, *** $P < 0.001$. See Figures S3 for all
 755 regression results.